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Process for preparing 1,1,1-trifluoro-2-chloroethane.

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The present invention relates to an improvement in the preparation of 1,1,1-trifluoro-2-chloroethane (hereinafter referred to as HCFC 133a), carried out by hydrofluorination of trichloroethylene (TCE) in the gas phase, in the presence of chrome oxide and/or chrome oxyfluorides or fluorides as catalysts.

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Preparations of HCFC 133a according to such method are broadly described in the art; in this connection, reference should be made to patents: US 3,752,850; 2,637,747; 2,885,427; 3,755,477; GB 1,000,485; DE 1,104,496.

A serious drawback affecting the preparation of HCFC 133a according to the abovesaid method is represented by the decay of the catalyst activity during the reaction.

So, for example, if a mixture of trichloroethylene and anhydrous HF in a 1/6 molar ratio is conveyed onto a catalyst based on chrome oxide and/or chrome oxyfluoride or fluoride carried on fluorinated alumina, with a contact time of 5 seconds, the starting conversion of trichloroethylene is very high (typically of 90-95%), which, however, gradually decreases at a rate which is depending on temperature. For example, at 300 °C the conversion begins to decrease after 25 hours and sinks to 80% within 90 hours, while at 350 °C the conversion sinks to 70-75% in 50 hours.

To overcome this drawback it was suggested to add air or oxygen to the reagent mixture.

Such method, however, is affected by the serious drawback of promoting dismutation reactions of the reaction intermediate products, giving rise to great amounts of undesired products. For example, maintaining the abovesaid conditions, with the exception of the addition of air in a O₂/trichloroethylene molar ratio equal to 1/6, about 6% by weight of dismutation products form, while in the absence of air such products form only in amounts of 0.5-0.6% by weight.

Furthermore, if it is operated in the presence of oxygen, the amount of 1,1-difluoro-2-chloroethylene (undesired product owing to its toxicity) rises from 0.02-0.03% to 0.25-0.30% by weight.

Lastly, operating in the presence of oxygen, the trichloroethylene conversion is slightly lower (88-90%) as compared with the cases when it is operated in the absence of oxygen.

It has now surprisingly been found by the Applicant that it is possible to maintain for a long period of time the catalyst activity and selectivity by adding to the reaction mixture, composed of trichloroethylene and of hydrofluoric acid, little amounts of HCFC 133a, what, in terms of plant engineering, is equivalent to recycle to the reactor a little part of the reaction product, optionally comprising the unreacted trichloroethylene and hydrofluoric acid.

A further advantage of this operating method resides in the fact that the formation of by-products decreases instead of increasing, as conversely happens when it is operated in the presence of oxygen.

The amount of HCFC 133a to be added to the reagent mixture sent to the reactor shall be at least of 7 mols %, and preferably of at least 10 mols %, referred to the mixture with trichloroethylene.

Generally, HCFC 133a amounts ranging from 7 to 25 mols % and preferably from 10 to 20 mols %, calculated on its mixture with trichloroethylene, can be considered as suitable for the purposes of the invention.

The reaction temperature suitably ranges from 250 ° to 330 ° C, preferably from 280 ° to 300 ° C.

From European patent application 449,614 it is known a process for preparing 1,1,1-trifluoro-2-fluoroethane by hydrofluorination of trichloroethylene, which process comprises a step of preparing 1,1,1-trifluoro-2-chloroethane (step D) via hydrofluorination of mixtures of said 1,1,1-trifluoro-2-chloroethane with trichloroethylene.

In such step, however, the amount of 1,1,1-trifluoro-2-chloroethane in the mixture is always by far prevailing on the amount of trichloroethylene (which, according to the examples, is of about 15 mols %), wherefore the conditions of the present invention do not exist in such process, nor the advantages are inferable therefrom.

Thus, it is an object of the present invention to provide a process for preparing HCFC 133a by hydrofluorination of trichloroethylene with hydrofluoric acid in the gas phase, in the presence of catalysts composed of chrome oxide and/or chrome oxyfluorides or fluorides and carried on at least partially fluorinated alumina, carried out under the above-described conditions.

The catalyst can be prepared according to any method of the art.

Preferably it is prepared by impregnating an alumina carrier, which had been at least partially previously fluorinated, with an aqueous solution of a trivalent chrome salt, by drying the whole and activating with air or nitrogen, either or not in the presence of water and/or crystallization water which can act as an oxidizer, at temperatures ranging fromn 200 ° to 600 °C, but preferably from 350 °C to 500 °C, in order to convert chrome into Cr₂O₃.

Preferably, but not necessarily, the catalyst is then treated, prior to the use, with gaseous HF, at a temperature from 200° to 450°C, optionally in the same reactor to be used for the reaction of trichloroethylene and HF.

By at least partially fluorinated alumina is meant herein the alumina comprising at least 50%, but preferably at least 90% by weight of AIF₃.

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Such a fluorinated alumina is preparable by hydrofluorination of alumina according to what is described in FR patent No. 1,383,927.

The chrome amount on the carrier preferably ranges from 1 to 15% by weight calculated on the total weight of the catalyst.

The carrier can be in the form of a powder having particle size from 20 to 200 microns, or in the form of pellets.

The at least partially fluorinated alumina used as a carrier comprises AIF₃ mainly in the gamma and/or beta forms; however also the delta form can be present up to amounts of 30% by weight.

Also AIF₃ in the alpha form can be present, however in little amounts.

Another method of preparing the catalyst comprises soaking the alumina with an aqueous solution of a chrome salt, drying and then, optionally subjecting the resulting product to fluorination with HF till obtaining a fluorination of the alumina for at least 50% and preferably for at least 90%.

A further method comprises coprecipitating the aluminium and chrome hydroxides, drying them, subjecting them to calcination in order to convert them into mixed oxides and, lastly, treating them with HF till obtaining the desired alumina fluorination degree.

In the trichloroethylene hydrofluorination reaction it is preferably operated with HF/trichloroethylene molar ratios higher than 3/1. Generally, ratios between 5/1 and 10/1 are utilized.

The pressure is not critical; generally it is operated at atmospheric pressure.

The following examples are given to illustrate the present invention, without being however a limitation thereof.

EXAMPLE 1

A catalyst was prepared by impregnating fluorinated alumina (containing 95% by weight of AlF₃) with an aqueous solution of $CrCl_3.6H_2O$, by drying and activating then the catalyst by means of treatment with N_2 at $400 \, ^{\circ}C$ for 8-10 hours.

390 cc of such catalyst were introduced into a reactor consisting of an INCONEL^(R) tube having a diameter of 50 mm, electrically heated and equipped with a sintered INCONEL fritted bottom.

Then, at a temperature of 300 °C there were fed 0.686 mols/h of trichloroethylene, 0.171 mols/h of HCFC 133a and 5.14 mols/h of anhydrous hydrofluoric acid, thereby obtaining a contact time of 5 seconds (measured as the ratio of the reagent volume at the reaction temperature to the volume of the catalytic bed at rest), a HF/organic products ratio of 6/1 and a HCFC 133a/trichloroethylene ratio of 1/4.

The gases leaving the reactor were scrubbed in water to eliminate their acidity, were dried and analyzed via GLC.

The following analysis, expressed in mols per cent, carried out after 48.5 hours, is representative: Al34a = 1.9; A133a = 92.3; TCE = 4.8; A1121 = 0.9; A143a = 0.1; besides negligible amounts of other products.

The trichloroethylene conversion was of 93.8% and after more than 70 hours of run under these conditions it was still the same, without any sign of decay of the catalyst performances, as well as the selectivity for 133a.

15 EXAMPLE 2

It was operated in the same reactor and under the same test conditions of example 1, varying only the 133a/trichloroethylene molar ratio, which in this case was equal to 1/9.

The catalytic bed volume was slightly smaller (350 cc) and therefore the flow rates were reduced in order to keep the contact time constant. The other operative conditions were as follows: HF = 3.94 mols/h; HCFC 133a = 0.066; TCE = 0.59.

Operating as in example 1, rather constant results in the course of time were obtained. The following analysis of the effluents, carried out after 63 hours, is representative:

A134a = 1.5; A133a = 90.7; TCE = 6.5; A1121 = 0.9; A1122 = 0.1; other by-products = 0.2.

The trichloroethylene conversion was equal to 92.6% and during a run of more than 85 hours it remained at 90-95% without signs of catalyst exhaustion.

EXAMPLE 3 (comparative)

it was operated in the same reactor and under the same test conditions of the preceding examples. 390 g, equal to 300 cc, of a previously activated catalyst, as is described in example 1, were charged, and at 300°C there were fed 4.10 mols/h of anhydrous hydrofluoric acid and 0.68 mols/h of trichloroethylene, so obtaining a contact time of 5 seconds and a HF/TCE ratio equal to 6.

The starting conversion was of 95-96% and gradually decreased to about 70% during a 68-hour run.

The reaction products were substantially the same as in the preceding examples as regards both nature and distribution, the conversion being equal; the following per cent analysis, carried out after a 20-hour run, is representative of obtained products: A134a = 1.3; A133a = 92.4; TCE = 4.4; A1121 = 1.1; A1122 = negligible; other by-products = 0.6.

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EXAMPLE 4 (use of oxygen)

390 g, equal to 300 cc, of catalyst already activated according to example 1 were charged into the reactor of the preceding examples. At 350 °C there were fed 0.21 mols/h of trichloroethylene, 1.73 mols/h of anhydrous HF and 4.15 l/h of air (equal to 0.035 mols of O_2), so obtaining a contact time of 10 seconds, a HF/trichloroethylene molar ratio of about 8/1 and a O_2 /trichloroethylene molar ratio equal to 1/6.

Operating as in the preceding examples there were obtained the results represented by the following mols per cent analysis of the products flowing from the reactor: A134a = 2.3; A133a = 81.3; TCE = 8.9; intermediates = 2.0; A1122 = 0.4; other by-products = 5.0.

The trichloroethylene conversion was equal to 91.1% and the by-products amounted to about 5.5% of the converted product.

Claims

- A process for preparing 1,1,1-trifluoro-2chloroethane by reacting, in the gas phase, trichloroethylene with HF in the presence of a catalyst composed of chrome trioxide and/or oxyfluoride and/or fluoride carried on AIF₃, characterized in that trichloroethylene in admixture with 7-25 mols % of 1,1,1-trifluoro-2chloroethane is fed to the hydrofluorination reactor.
- The process of claim 1, wherein 1,1,1-trifluoro-2-chloroethane in admixture with trichloroethylene is in an amount ranging from 10 to 20 mols % referred to the total of said mixture.
- The process of claims 1 and 2, wherein the reaction is caused to occur at a temperature ranging from 250 to 330 °C.

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EUROPEAN SEARCH REPORT

Application Number

EP 93 11 2630

DOCUMENTS CONSIDERED TO BE RELEVANT				
Category	Citation of document with of relevant p	indication, where appropriate, assages	Relevant to claim	CLASSIFICATION OF THE APPLICATION (Int. CL5)
Y	EP-A-0 446 869 (DA * page 3, column 3 38; claim 1 *	IKIN INDUSTRIES LIMITED), line 6 - page 7, line	1-3	C07C17/20 C07C19/08
Y	EP-A-0 408 005 (AUS* the whole document		1-3	
A	WO-A-9 008 755 (E.I AND COMPANY) * page 11 - page 15	I. DU PONT DE NEMOURS	1-3	
A,D	EP-A-0 449 614 (IMF INDUSTRIES PLC) * the whole documen	•	1-3	
A,D	US-A-2 885 427 (ROE * examples 1,2 *	BERT P. RUH ET AL.)	1	
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				TECHNICAL FIELDS SEARCHED (Int. Cl.5)
				C07C
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	The present search report has b	een drawn up for all claims		
	Place of search	Date of completion of the search		Economic
BERLIN 30 N		30 NOVEMBER 1993		RUFET J.
X : partic Y : partic docum A : techn	ATEGORY OF CITED DOCUMES cularly relevant if taken alone cularly relevant if combined with and ment of the same category tological background written disclosure	E : earlier parent docu after the filing dat	ment, but publi te the application other reasons	shed on, or

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